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L7: Entry 1 of 2

File: USPT

Feb 1, 2000

US-PAT-NO: 6020288

DOCUMENT-IDENTIFIER: US 6020288 A

TITLE: Methods and compositions for enhancing cytochrome P450 in plants

DATE-ISSUED: February 1, 2000

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Nonomura; Arthur M.	Boxborough	MA	01719	N/A
Benson; Andrew A.	La Jolla	CA	92037	N/A
Nishio; John N.	Laramie	WY	82070-3917	N/A

APPL-NO: 8 / 927415

DATE FILED: September 11, 1997

## PARENT-CASE:

RELATED APPLICATIONS This application is a continuation-in-part of U.S. patent application Ser. No. 08/610,928, filed Mar. 5, 1996, now U.S. Pat. No. 5,846,908, which is a continuation-in part of U.S. patent application Ser. No. 08/399,399, filed Mar. 6, 1995; which was a continuation-in part of Ser. No. 08/351,348 filed Dec. 9, 1994, now U.S. Pat. No. 5,597,400 issued on Jan. 28, 1997, which was a continuation-in-part of U.S. patent application Ser. No. 07/901,366, filed on Jun. 19, 1992. The full disclosures of each of these patent applications are incorporated herein by reference. Related international application are PCT/US96/02444 (equivalent of Ser. No. 08/610,928) and PCT/US93/05676 (equivalent of Ser. No. 08/351,348).

INT-CL: [6] A01N 31/00, A01N 37/00, A01N 43/22, A01N 57/02

US-CL-ISSUED: 504/127, 504/128, 504/130, 504/136, 504/138, 504/140, 504/142, 504/143, 504/144, 504/149

US-CL-CURRENT: 504/127, 504/128, 504/130, 504/136, 504/138, 504/140, 504/142, 504/143, 504/144, 504/149

FIELD-OF-SEARCH: 504/127, 504/128, 504/130, 504/136, 504/138, 504/140, 504/142, 504/143, 504/144, 504/149

## PRIOR-ART-DISCLOSED:

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PAT-NO	ISSUE-DATE	PATENTEE-NAME	US-CL
<u>3897241</u>	July 1975	Washio et al.	71/113
<u>4799953</u>	January 1989	Danzig et al.	71/98
<u>4846877</u>	July 1989	Azuma et al.	71/92
<u>5298482</u>	March 1994	Tanaka et al.	504/320
<u>5300540</u>	April 1994	Masters	523/309
<u>5532204</u>	July 1996	Joshi	504/118
<u>5597400</u>	January 1997	Nonomura et al.	71/28

## FOREIGN PATENT DOCUMENTS

FOREIGN-PAT-NO	PUBN-DATE	COUNTRY	US-CL
0 465 907 A1	January 1992	EPX	
2689905	October 1993	FRX	
2004856	April 1979	GBX	

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- cytochromes P450" *Nature Biotechnol.* 15:784-788 (1997).
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ART-UNIT: 166

PRIMARY-EXAMINER: Clardy; S. Mark

ATTY-AGENT-FIRM: Nields, Lemack &amp; Dingman

## ABSTRACT:

The present invention provides methods for treating plants which comprise application of an oxidant that induces NADPH:cytochrome P450 reductase and application of a reductant that induces cytochrome P450 monooxygenase. The present invention also provides methods for increasing cytochrome P450 in plants and for enhancing the growth of plants. The present invention also provides compositions and systems useful in the methods of the present invention.

38 Claims, 1 Drawing figures

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw Desc	Image
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 2. Document ID: US 5254466 A

L7: Entry 2 of 2

File: USPT

Oct 19, 1993

US-PAT-NO: 5254466

DOCUMENT-IDENTIFIER: US 5254466 A

TITLE: Site-specific modification of the candida tropicales genome

DATE-ISSUED: October 19, 1993

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Picataggio; Stephen	Santa Rosa	CA	N/A	N/A
Deanda; Kristine	Gronon	CA	N/A	N/A
Eirich; L. Dudley	Santa Rosa	CA	N/A	N/A

## ASSIGNEE-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY	TYPE	CODE
Henkel Research Corporation	Santa Rosa	CA	N/A	N/A	02	

APPL-NO: 7/ 432091

DATE FILED: November 6, 1989

INT-CL: [5] C12N 15/09, C12P 7/44  
 US-CL-ISSUED: 435/142; 435/172.1, 435/172.3, 435/924, 435/254.22  
 US-CL-CURRENT: 435/142; 435/254.22, 435/477, 435/481, 435/483, 435/484,  
435/490, 435/6, 435/924  
 FIELD-OF-SEARCH: 435/67.1, 435/142, 435/172.3, 435/172.1, 435/940, 435/255,  
 435/924, 935/22, 935/28

PRIOR-ART-DISCLOSED:

U.S. PATENT DOCUMENTS

PAT-NO	ISSUE-DATE	PATENTEE-NAME	US-CL
<u>4735901</u>	April 1988	Kurtz et al.	435/172.3

FOREIGN PATENT DOCUMENTS

FOREIGN-PAT-NO	PUBN-DATE	COUNTRY	US-CL
183070	April 1986	EPX	

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Proceeding National Academy of Science, vol. 76 No. 10 issued Oct. 1979. S. Scherer et al. "Replacement of Chromosome Segments with altered DNA sequences constructed in vitro". pp. 4951-4955.

ART-UNIT: 185

PRIMARY-EXAMINER: Schwartz; Richard A.

ASSISTANT-EXAMINER: LeGuyader; J.

ATTY-AGENT-FIRM: Szoke; Ernest G. Jaeschke; Wayne C. Drach; John E.

ABSTRACT:

The POX genes of *C. tropicalis* are disrupted resulting in the complete blockage of the beta-oxidation pathway in the strain. Fermentation of *C. tropicalis* cells having disrupted genes on alkane, fatty acid and fatty acid ester substrates produces substantially pure dicarboxylic acids in substantially quantitative yield.

30 Claims, 5 Drawing figures

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L9: Entry 1 of 2

File: USPT

Feb 1, 2000

US-PAT-NO: 6020288

DOCUMENT-IDENTIFIER: US 6020288 A

TITLE: Methods and compositions for enhancing cytochrome P450 in plants

DATE-ISSUED: February 1, 2000

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Nonomura; Arthur M.	Boxborough	MA	01719	N/A
Benson; Andrew A.	La Jolla	CA	92037	N/A
Nishio; John N.	Laramie	WY	82070-3917	N/A

US-CL-CURRENT: 504/127; 504/128, 504/130, 504/136, 504/138, 504/140, 504/142,  
504/143, 504/144, 504/149[Full](#) | [Title](#) | [Citation](#) | [Front](#) | [Review](#) | [Classification](#) | [Date](#) | [Reference](#) | [Claims](#) | [KMIC](#) | [Drawn Desc](#) | [Image](#) **2. Document ID: US 5254466 A**

L9: Entry 2 of 2

File: USPT

Oct 19, 1993

US-PAT-NO: 5254466

DOCUMENT-IDENTIFIER: US 5254466 A

TITLE: Site-specific modification of the candida tropica genome

DATE-ISSUED: October 19, 1993

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Picataggio; Stephen	Santa Rosa	CA	N/A	N/A
Deanda; Kristine	Graton	CA	N/A	N/A
Eirich; L. Dudley	Santa Rosa	CA	N/A	N/A

US-CL-CURRENT: 435/142; 435/254.22, 435/477, 435/481, 435/483, 435/484,  
435/490, 435/6, 435/924[Full](#) | [Title](#) | [Citation](#) | [Front](#) | [Review](#) | [Classification](#) | [Date](#) | [Reference](#) | [Claims](#) | [KMIC](#) | [Drawn Desc](#) | [Image](#)

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**WEST****Generate Collection****Search Results - Record(s) 1 through 3 of 3 returned.** **1. Document ID: US 6174673 B1**

L17: Entry 1 of 3

File: USPT

Jan 16, 2001

US-PAT-NO: 6174673

DOCUMENT-IDENTIFIER: US 6174673 B1

TITLE: High throughput screening for novel enzymes

DATE-ISSUED: January 16, 2001

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Short; Jay M.	Encinitas	CA	N/A	N/A
Keller; Martin	San Diego	CA	N/A	N/A

## ASSIGNEE-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY	TYPE CODE
Diversa Corporation	San Diego	CA	N/A	N/A	02

APPL-NO: 9/ 098206

DATE FILED: June 16, 1998

## PARENT-CASE:

This application is a continuation-in-part of U.S. application Ser. No. 08/876,276, filed Jun. 16, 1997.

INT-CL: [7] C12Q 1/68

US-CL-ISSUED: 435/6; 435/69.1, 435/440, 435/471, 435/476, 435/320.1

US-CL-CURRENT: 435/6; 435/320.1, 435/440, 435/471, 435/476, 435/69.1FIELD-OF-SEARCH: 435/6, 435/69.1, 435/440, 435/320.1, 435/471, 435/476

## PRIOR-ART-DISCLOSED:

## U. S. PATENT DOCUMENTS

PAT-NO	ISSUE-DATE	PATENTEE-NAME	US-CL
<u>4399219</u>	August 1983	Weaver	435/34
<u>4643968</u>	February 1987	Weaver	435/32
<u>4647536</u>	March 1987	Mosbach et al.	435/177
<u>4916060</u>	April 1990	Weaver	435/29
<u>4959301</u>	September 1990	Weaver et al.	435/5
<u>5055390</u>	October 1991	Weaver et al.	435/5
<u>5225332</u>	July 1993	Weaver et al.	435/29
<u>5824485</u>	October 1998	Thompson et al.	435/6

## FOREIGN PATENT DOCUMENTS

FOREIGN-PAT-NO	PUBN-DATE	COUNTRY	US-CL
WO 98/56904	December 1998	WOX	
WO 99/49315	September 1999	WOX	
WO99/54494	October 1999	WOX	

## OTHER PUBLICATIONS

Zhang et al. FASEB J. vol. 5, pp. 3108-3113, 1991.  
 Plovins et al. Applied and Environmental Microbiology. vol. 60(12), pp. 4638-4641, 1994.

ART-UNIT: 166

PRIMARY-EXAMINER: Yucel; Remy

ATTY-AGENT-FIRM: Gray Gary Ware &amp; Freidenrich LLP Haile; Lisa A.

## ABSTRACT:

Disclosed is a process for identifying clones having a specified activity of interest, which process comprises (i) generating one or more expression libraries derived from nuclei acid directly isolated from the environment; and (iii) screening said libraries utilizing a fluorescence activated cell sorter to identify said clones. More particularly, this is a process for identifying clones having a specified activity of interest by (i) generating one or more expression libraries derived from nucleic acid directly or indirectly isolated from the environment; (ii) exposing said libraries to a particular substrate or substrates of interest; and (iii) screening said exposed libraries utilizing a fluorescence activated cell sorter to identify clones which react with the substrate or substrates. Also provided is a process for identifying clones having a specified activity of interest by (i) generating one or more expression libraries derived from nucleic acid directly or indirectly isolated from the environment; and (ii) screening said exposed libraries utilizing an assay requiring co-encapsulation, a binding event or the covalent modification of a target, and a fluorescence activated cell sorter to identify positive clones.

23 Claims, 18 Drawing figures

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KMC	Drawn Desc	Image
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 2. Document ID: US 6168919 B1

L17: Entry 2 of 3

File: USPT

Jan 2, 2001

US-PAT-NO: 6168919

DOCUMENT-IDENTIFIER: US 6168919 B1

TITLE: Screening methods for enzymes and enzyme kits

DATE-ISSUED: January 2, 2001

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Short; Jay M.	Encinitas	CA	N/A	N/A

## ASSIGNEE-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY	TYPE CODE
Diversa Corporation	San Diego	CA	N/A	N/A	02

APPL-NO: 8 / 983367

DATE FILED: September 30, 1998

PCT-DATA:

APPL-NO	DATE-FILED	PUB-NO	PUB-DATE	371-DATE	102(E)-DATE
PCT/US96/11854	July 17, 1996	WO97/04077	Feb 6, 1997	Sep 30, 1998	Sep 30, 1998

INT-CL: [7] C12Q 1/68, C12Q 1/00, C12P 19/34, C12N 15/64  
 US-CL-ISSUED: 435/6; 435/4, 435/91.1, 435/91.4, 435/91.41, 435/252.3, 435/183,  
 435/320.1, 435/325, 536/23.1, 536/23.2, 536/23.4  
 US-CL-CURRENT: 435/6; 435/183, 435/252.3, 435/320.1, 435/325, 435/4, 435/91.1,  
 435/91.4, 435/91.41, 536/23.1, 536/23.2, 536/23.4  
 FIELD-OF-SEARCH: 435/4, 435/6, 435/91.1, 435/91.4, 435/91.41, 435/252.3,  
 435/183, 435/320.1, 435/325, 536/23.1, 536/23.2, 536/23.4

PRIOR-ART-DISCLOSED:

U.S. PATENT DOCUMENTS

PAT-NO	ISSUE-DATE	PATENTEE-NAME	US-CL
<u>5171684</u>	December 1992	Yen et al.	435/252.3
<u>5712146</u>	January 1998	Khosla et al.	435/252.35
<u>5958672</u>	September 1999	Short	435/4

OTHER PUBLICATIONS

Lactic Dehydrogenase, Sigma catalog, p. 634, 1997.  
 Anderson et al., Met. Enzymol., vol. 68, pp. 428-436, 1979.  
 Promega Catalog, p. 205, 1993.  
 Sambrook et al., Molecular Cloning, Cold Spring Harbor Laboratory Press, vol. 2, p. 9.30 and pp. 12.1-12.20, 1989.\*

ART-UNIT: 162

PRIMARY-EXAMINER: Achutamurthy; Ponnathapu

ASSISTANT-EXAMINER: Tung; Peter P.

ATTY-AGENT-FIRM: Gray, Cary, Ware & Freidenrich LLP Haile; Lisa A.

ABSTRACT:

Recombinant enzyme libraries and kits where a plurality of enzymes are each characterized by different physical and/or chemical characteristics and classified by common characteristics. The characteristics are determined by screening of recombinant enzymes expressed by a DNA library produced from various microorganisms. Also disclosed is a process for identifying clones of a recombinant library which express a protein with a desired activity by screening a library of expression clones randomly produced from DNA of at least one microorganism, said screening being effected on expression products of said clones to thereby identify clones which express a protein with a desired activity. Also disclosed is a process of screening clones having DNA from an uncultivated microorganism for a specified protein activity by screening for a specified protein activity in a library of clones prepared by (I) recovering DNA from a DNA population derived from at least one uncultivated microorganism; and (ii) transforming a host with recovered DNA to produce a library of clones which is screened for the specified protein activity.

9 Claims, 8 Drawing figures

<a href="#">Full</a>	<a href="#">Title</a>	<a href="#">Citation</a>	<a href="#">Front</a>	<a href="#">Review</a>	<a href="#">Classification</a>	<a href="#">Date</a>	<a href="#">Reference</a>	<a href="#">Claims</a>	<a href="#">KMC</a>	<a href="#">Drawn Desc</a>	<a href="#">Image</a>
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 3. Document ID: US 5254466 A

L17: Entry 3 of 3

File: USPT

Oct 19, 1993

US-PAT-NO: 5254466

DOCUMENT-IDENTIFIER: US 5254466 A

TITLE: Site-specific modification of the candida tropicales genome

DATE-ISSUED: October 19, 1993

## INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Picataggio; Stephen	Santa Rosa	CA	N/A	N/A
Deanda; Kristine	Graton	CA	N/A	N/A
Eirich; L. Dudley	Santa Rosa	CA	N/A	N/A

## ASSIGNEE-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY	TYPE CODE
Henkel Research Corporation	Santa Rosa	CA	N/A	N/A	02

APPL-NO: 7/ 432091

DATE FILED: November 6, 1989

INT-CL: [5] C12N 15/09, C12P 7/44

US-CL-ISSUED: 435/142; 435/172.1, 435/172.3, 435/924, 435/254.22

US-CL-CURRENT: 435/142; 435/254.22, 435/477, 435/481, 435/483, 435/484,  
435/490, 435/6, 435/924FIELD-OF-SEARCH: 435/67.1, 435/142, 435/172.3, 435/172.1, 435/940, 435/255,  
435/924, 935/22, 935/28

## PRIOR-ART-DISCLOSED:

## U.S. PATENT DOCUMENTS

PAT-NO	ISSUE-DATE	PATENTEE-NAME	US-CL
<u>4735901</u>	April 1988	Kurtz et al.	435/172.3

## FOREIGN PATENT DOCUMENTS

FOREIGN-PAT-NO	PUBN-DATE	COUNTRY	US-CL
183070	April 1986	EPX	

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"Export of the carboxy-terminal Portion of Acyl-CoA oxidase into Peroxisomes of *Candida tropicalis*" pp. 217-252.

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Journal of Bacteriology, vol. 172, No. 3 issue Aug. 1990, [O. C. Haas et al., "Development of an integrative DNA Transformation System for the yeast *Candida tropicalis*", pp. 4571-4577.

Gene vol. 51, issued 1987, W. W. Murray et al, "The primary structure of a peroxisomal fatty acyl-CoA oxidase from the yeast Canada tropicalis p K233", pp. 119-128.

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ART-UNIT: 185

PRIMARY-EXAMINER: Schwartz; Richard A.

ASSISTANT-EXAMINER: LeGuyader; J.

ATTY-AGENT-FIRM: Szoke; Ernest G. Jaeschke; Wayne C. Drach; John E.

#### ABSTRACT:

The POX genes of *C. tropicalis* are disrupted resulting in the complete blockage of the beta-oxidation pathway in the strain. Fermentation of *C. tropicalis* cells having disrupted genes on alkane, fatty acid and fatty acid ester substrates produces substantially pure dicarboxylic acids in substantially quantitative yield.

30 Claims, 5 Drawing figures

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KMC	Draw. Desc	Image
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L2 1 SEA ABB=ON PLU=ON 9038-14-6/RN  
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L3 1 SEA ABB=ON PLU=ON 9023-03-4/RN  
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L4 SEL PLU=ON L2 1- CHEM : 26 TERMS

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SET SMARTSELECT ON

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SET SMARTSELECT OFF

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L7 7478 SEA ABB=ON PLU=ON L5

L8 1165 SEA ABB=ON PLU=ON L6 AND L7

L9 188643 SEA ABB=ON PLU=ON DICARBOXYLIC ACID# OR (CARBOXYLIC ACIDS  
(L) DICARBOXYLIC) OR CARBOXYLIC ACID# OR MONOCARBOXYLIC ACID#

L10 0 SEA ABB=ON PLU=ON L8 (L) L9

L11 5883 SEA ABB=ON PLU=ON (HYDROXYLATION (L) .OMEGA.-HYDROXYLATION)  
OR (HYDROXYLATION (L) ALKANE#) OR (.ALPHA.-HYDROXYLATION) OR  
(HYDROXYLATION (L) .ALPHA.)

L12 0 SEA ABB=ON PLU=ON L9 AND L10

L13 149 SEA ABB=ON PLU=ON L9 AND L11

L14 8 SEA ABB=ON PLU=ON L13 AND (L6 OR L7)

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L16 8 SEA L15

L17 5 SEA ABB=ON PLU=ON L16 AND (PREP/RL OR PREPAR? OR SYNTH? OR  
MAK?)  
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FILE CAPLUS

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L17 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 2000:133824 CAPLUS  
DOCUMENT NUMBER: 132:162018  
TITLE: DNA shuffling of **monooxygenase** genes for  
production of industrial chemicals  
INVENTOR(S): Affholter, Joseph A.; Davis, Christopher; Selifonov,  
Sergey A.  
PATENT ASSIGNEE(S): Maxygen, Inc., USA  
SOURCE: PCT Int. Appl., 153 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000009682	A1	20000224	WO 1999-US18424	19990812
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
AU 9953479	A1	20000306	AU 1999-53479	19990812
PRIORITY APPLN. INFO.:			US 1998-96271	P 19980812
			US 1999-130810	P 19990423
			WO 1999-US18424	W 19990812

OTHER SOURCE(S): MARPAT 132:162018  
AB This invention provides improved **monooxygenases**, dehydrogenases, and transferases that are useful for the biocatalytic **synthesis** of compds. such as .alpha.-hydroxycarboxylic acids, and aryl- and alkyl-, hydroxy compds. The polypeptides provided herein are improved in properties such as regioselectivity, enzymic activity, stereospecificity, and the like. Methods for obtaining recombinant polynucleotides that encode these improved polypeptides are also provided, as are organisms that express the polypeptides and are thus useful for carrying out said biocatalytic **syntheses**. In the methods for obtaining **monooxygenase** genes, a plurality of parental forms (homologs) of a selected nucleic acid are recombined. The selected nucleic acid derived either from one or more parental nucleic acid(s) which encodes a **monooxygenase** enzyme, or a fragment thereof, or from a parental nucleic acid which does not encode **monooxygenase**, but which is a candidate for DNA shuffling to develop **monooxygenase** activity. The plurality of forms of the selected nucleic acid differ from each other in at lease one (and typically two or more) nucleotides, and, upon recombination, provide a library of recombinant **monooxygenase** nucleic acids. The library can be an *in vitro* set of mols., or present in cells, phage or the like. The library is screened to identify at least one recombinant **monooxygenase** nucleic acid that exhibits distinct or improved **monooxygenase** activity compared to the parental nucleic acid or nucleic acids. Also provided by the invention

are methods for increasing said solvent resistance of organisms that are used in the **synthetic** methods.

REFERENCE COUNT:

18

REFERENCE(S):

- (1) Affymax Tech Nv; WO 9720078 A 1997 CAPLUS
- (2) Agency Of Ind Sci & Technology; JP 05-049474 A 1993 CAPLUS
- (3) Aoyama, T; JOURNAL OF BIOLOGICAL CHEMISTRY 1989, V264(18), P10388 CAPLUS
- (4) Crameri, A; NATURE 1998, V391, P288 CAPLUS
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ALL CITATIONS AVAILABLE IN THE RE FORMAT

- TI DNA shuffling of **monooxygenase** genes for production of industrial chemicals
- AB This invention provides improved **monooxygenases**, dehydrogenases, and transferases that are useful for the biocatalytic **synthesis** of compds. such as .alpha.-hydroxycarboxylic acids, and aryl- and alkyl-, hydroxy compds. The polypeptides provided herein are improved in properties such as regioselectivity, enzymic activity, stereospecificity, and the like. Methods for obtaining recombinant polynucleotides that encode these improved polypeptides are also provided, as are organisms that express the polypeptides and are thus useful for carrying out said biocatalytic **syntheses**. In the methods for obtaining **monooxygenase** genes, a plurality of parental forms (homologs) of a selected nucleic acid are recombined. The selected nucleic acid derived either from one or more parental nucleic acid(s) which encodes a **monooxygenase** enzyme, or a fragment thereof, or from a parental nucleic acid which does not encode **monooxygenase**, but which is a candidate for DNA shuffling to develop **monooxygenase** activity. The plurality of forms of the selected nucleic acid differ from each other in at lease one (and typically two or more) nucleotides, and, upon recombination, provide a library of recombinant **monooxygenase** nucleic acids. The library can be an in vitro set of mols., or present in cells, phage or the like. The library is screened to identity at least one recombinant **monooxygenase** nucleic acid that exhibits distinct or improved **monooxygenase** activity compared to the parental nucleic acid or nucleic acids. Also provided by the invention are methods for increasing said solvent resistance of organisms that are used in the **synthetic** methods.
- ST **monooxygenase** industrial **synthesis** DNA shuffling gene
- IT Bioreactors
- Nucleic acid library
- Phage display library
- Recombination, genetic
- Regiochemistry  
(DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)
- IT Gene
- RL: BPR (Biological process); BUU (Biological use, unclassified); BIOL (Biological study); PROC (Process); USES (Uses)  
(DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)
- IT Amines, biological studies
- RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study); PROC (Process)  
(N-dealkylation; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)
- IT Ethers, biological studies

RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(O-dealkylation; DNA shuffling of **monooxygenase** genes for  
prodn. of industrial chems.)

IT Thiols (organic), biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(S-dealkylation; DNA shuffling of **monooxygenase** genes for  
prodn. of industrial chems.)

IT Ethers, biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(allyl, substrate; DNA shuffling of **monooxygenase** genes for  
prodn. of industrial chems.)

IT Baeyer-Villiger oxidation  
Dealkylation  
Decarboxylation  
Dehalogenation  
Dehydrogenation  
Epoxidation  
Hydroxylation  
Oxidation  
(enzymic; DNA shuffling of **monooxygenase** genes for prodn. of  
industrial chems.)

IT Alkenes, biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(epoxidn.; DNA shuffling of **monooxygenase** genes for prodn. of  
industrial chems.)

IT Aromatic hydrocarbons, biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(halo, oxidative dehalogenation; DNA shuffling of **monooxygenase**  
genes for prodn. of industrial chems.)

IT Carboxylic acids, biological studies  
RL: BPN (Biosynthetic preparation); MFM (Metabolic formation); BIOL  
(Biological study); FORM (Formation, nonpreparative); PREP  
(Preparation)  
(hydroxy, conversion from olefins; DNA shuffling of  
**monooxygenase** genes for prodn. of industrial chems.)

IT Alkanes, biological studies  
Aromatic hydrocarbons, biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(hydroxylation; DNA shuffling of **monooxygenase**  
genes for prodn. of industrial chems.)

IT Chemicals  
(industrial; DNA shuffling of **monooxygenase** genes for prodn.  
of industrial chems.)

IT Solvents  
(org., resistance to; DNA shuffling of **monooxygenase** genes  
for prodn. of industrial chems.)

IT Sulfonylureas  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(oxygenation; DNA shuffling of **monooxygenase** genes for prodn.  
of industrial chems.)

IT Proteins, specific or class  
RL: BPN (Biosynthetic preparation); BIOL (Biological study); PREP  
(Preparation)

(solvent-resistant; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT Glycols, biological studies  
RL: BPN (Biosynthetic preparation); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); **PREP (Preparation)**  
(vicinal, conversion from olefins; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 9003-99-0P, Peroxidase 9013-18-7P, Acyl-CoA Ligase 9023-26-1P, Coenzyme A transferase 9024-04-8P, Mandelate racemase 9031-56-5P, Ligase 9033-07-2P, Glycosyltransferase 9033-25-4P, Methyltransferase 9035-51-2P, Cytochrome P 450, **preparation** 9035-82-9P, Dehydrogenase **9038-14-6P**, **Monooxygenase** 9047-61-4P, Transferase 9048-63-9P, Epoxide hydrolase 9054-54-0P, Acyltransferase 9080-22-2P, Racemase 102055-73-2P, Oxo acid decarboxylase  
RL: BPN (Biosynthetic preparation); CAT (Catalyst use); BIOL (Biological study); **PREP (Preparation)**; USES (Uses)  
(DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 60-12-8DP, 2-Phenylethanol, substituted 60-12-8P, 2-Phenylethanol 100-51-6DP, Benzyl alcohol, substituted 100-51-6P, Benzyl alcohol, biological studies 122-97-4DP, 3-Phenylpropanol, substituted 122-97-4P, 3-Phenylpropanol  
RL: BPN (Biosynthetic preparation); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); **PREP (Preparation)**  
(DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 145-13-1P, Pregnenolone  
RL: BPN (Biosynthetic preparation); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); **PREP (Preparation)**  
(conversion from cholesterol; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 81093-37-0P, Pravastatin  
RL: BPN (Biosynthetic preparation); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); **PREP (Preparation)**  
(conversion from mevastatin; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 57-88-5, Cholesterol, biological studies  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study); PROC (Process)  
(conversion to pregnenolone; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 119-84-6, 3,4-Dihydrocoumarin 73573-88-3, Mevastatin  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study); PROC (Process)  
(hydroxylation; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 59865-13-3, Cyclosporin 79217-60-0, Cyclosporin  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study); PROC (Process)  
(modification; DNA shuffling of **monooxygenase** genes for prodn. of industrial chems.)

IT 78-79-5, Isoprene, biological studies 100-42-5, Styrene, biological studies 106-99-0, Butadiene, biological studies 107-02-8, Acrolein, biological studies 557-40-4, Diallyl ether 1321-74-0, Divinylbenzene, biological studies 1321-74-0D, Divinylbenzene, substituted 1746-13-0, Allyl phenyl ether 1746-13-0D, Allyl phenyl ether, substituted

25168-07-4, Vinylcyclohexene 40356-67-0, Vinylnorbornene  
RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);  
PROC (Process)  
(substrate; DNA shuffling of **monooxygenase** genes for prodn.  
of industrial chems.)

L17 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1999:77701 CAPLUS  
DOCUMENT NUMBER: 130:138397  
TITLE: Yeasts with elevated cytochrome P450 levels and their  
use in the manufacture of monoterpenal and diterpenal  
aliphatic carboxylates from alkanes  
INVENTOR(S): Fallon, Robert D.; Payne, Mark S.; Picataggio,  
Stephen  
K.; Wu, Shijun  
PATENT ASSIGNEE(S): E.I. Du Pont De Nemours and Company, USA  
SOURCE: PCT Int. Appl., 58 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 1  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9904014	A2	19990128	WO 1998-US14935	19980720
WO 9904014	A3	19990520		
W: AU, CA, IL, JP, NZ, US				
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,				
PT, SE				
AU 9884982	A1	19990210	AU 1998-84982	19980720
EP 1003881	A2	20000531	EP 1998-935806	19980720
R: DE, DK, ES, FR, GB, IT, NL, SE, IE				
PRIORITY APPLN. INFO.:			US 1997-53215	P 19970721
			WO 1998-US14935	W 19980720

AB Strains of *Pichia pastoris* and *Candida maltosa* that have increased cytochrome P 450 activity or lack the .beta.-oxidn. pathway that can be used to convert c6-C22 alkanes to monoterpenal and diterpenal **carboxylic acids** are described. Expression constructs carrying cytochrome P 450 genes under the control of powerful inducible promoters (AOX1, PGK) were constructed by std. methods and introduced into

*P. pastoris* or *C. maltosa* to give novel or increased P 450 activities. Disruption of .beta.-oxidn. in *C. maltosa* was achieved by insertional inactivation of the POX4 gene. A strain of *C. maltosa* cultured in a complete medium that was supplemented with dodecane 20 g/L when glucose was almost completely depleted was able to generate dodecanedioic acid at 3.4 g/h for 51 h.

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3.4 g/h for 51 h.

ST Pichia alkane hydroxylation carboxylic dicarboxylic acid manuf; cytochrome P450 Candida Pichia alkane carboxylic acid fermn; Candida alkane hydroxylation carboxylic dicarboxylic acid manuf

IT Carboxylic acids, preparation  
Dicarboxylic acids  
RL: BMF (Bioindustrial manufacture); BIOL (Biological study); PREP (Preparation)  
(C6-22; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT Fermentation  
(carboxylic acids, with transgenic yeasts; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT Plasmid vectors  
(pLPA1T, alkane monooxygenase gene on, expression in Pichia of; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT Plasmid vectors  
(pSW84, alkane monooxygenase and cytochrome reductase genes on, expression in Candida of; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT Plasmid vectors  
(pSW87, alkane monooxygenase and cytochrome reductase genes on, expression in Candida of; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT 9038-14-6, Monooxygenase 9059-16-9, Alkane monooxygenase 106178-16-9, Fatty acid monooxygenase  
RL: BPR (Biological process); BUU (Biological use, unclassified); CAT (Catalyst use); BIOL (Biological study); PROC (Process); USES (Uses)  
(in alkane utilization by yeasts; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

IT 693-23-2P, Dodecanedioic acid  
RL: BMF (Bioindustrial manufacture); BIOL (Biological study); PREP (Preparation)  
(manuf. from dodecane of; yeasts with elevated cytochrome P 450 levels and their use in manuf. of monoterminal and diterminal aliph. carboxylates from alkanes)

L17 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1993:5685 CAPLUS  
DOCUMENT NUMBER: 118:5685  
TITLE: Metabolic engineering of Candida tropicalis for the production of long-chain dicarboxylic acids  
AUTHOR(S): Picataggio, Stephen; Rohrer, Tracy; Deanda, Kristine; Lanning, Dawn; Reynolds, Robert; Mielenz, Jonathan; Eirich, L. Dudley  
CORPORATE SOURCE: Microb. Technol. Dep., Cognis Inc., Santa Rosa, CA, 95407, USA  
SOURCE: Bio/Technology (1992), 10(8), 894-8  
CODEN: BTCHDA; ISSN: 0733-222X

- DOCUMENT TYPE: Journal  
LANGUAGE: English
- AB An industrial strain of the yeast *C. tropicalis* was engineered for the efficient prodn. of long-chain **dicarboxylic acids**, which are important raw materials for the chem. industry. By sequential disruption of the 4 genes encoding both isoenzymes of the acyl-CoA oxidase which catalyzes the first reaction in the .beta.-oxidn. pathway, **alkane** and fatty acid substrates were successfully redirected to the .omega.-oxidn. pathway. Consequently, the conversion efficiency and chem. selectivity of their terminal oxidn. to the corresponding **dicarboxylic acids** was improved to 100%. The specific productivity of the bioconversion was increased further by amplification of the **cytochrome P 450 monooxygenase** and NADPH-cytochrome reductase genes encoding the rate-limiting .omega.-hydroxylase in the .omega.-oxidn. pathway. The amplified strains demonstrated increased .omega.-hydroxylase activity and a 30% increase in productivity compared to the .beta.-oxidn.-blocked strain in fermns. The bioconversion is effective for the selective terminal oxidn. of both satd. and unsatd. linear aliph. substrates with C12-22 chain-lengths and also avoids the undesirable chain modifications assocd. with passage through the .beta.-oxidn. pathway, such as unsatn., **hydroxylation**, or chain shortening. It is now possible to efficiently produce a wide range of previously unavailable satd. and unsatd. **dicarboxylic acids** with a high degree of purity.
- TI Metabolic engineering of *Candida tropicalis* for the production of long-chain **dicarboxylic acids**
- AB An industrial strain of the yeast *C. tropicalis* was engineered for the efficient prodn. of long-chain **dicarboxylic acids**, which are important raw materials for the chem. industry. By sequential disruption of the 4 genes encoding both isoenzymes of the acyl-CoA oxidase which catalyzes the first reaction in the .beta.-oxidn. pathway, **alkane** and fatty acid substrates were successfully redirected to the .omega.-oxidn. pathway. Consequently, the conversion efficiency and chem. selectivity of their terminal oxidn. to the corresponding **dicarboxylic acids** was improved to 100%. The specific productivity of the bioconversion was increased further by amplification of the **cytochrome P 450 monooxygenase** and NADPH-cytochrome reductase genes encoding the rate-limiting .omega.-hydroxylase in the .omega.-oxidn. pathway. The amplified strains demonstrated increased .omega.-hydroxylase activity and a 30% increase in productivity compared to the .beta.-oxidn.-blocked strain in fermns. The bioconversion is effective for the selective terminal oxidn. of both satd. and unsatd. linear aliph. substrates with C12-22 chain-lengths and also avoids the undesirable chain modifications assocd. with passage through the .beta.-oxidn. pathway, such as unsatn., **hydroxylation**, or chain shortening. It is now possible to efficiently produce a wide range of previously unavailable satd. and unsatd. **dicarboxylic acids** with a high degree of purity.
- ST genetic engineering *Candida* **dicarboxylic acid** prodn
- IT *Candida tropicalis*  
(genetic engineering of, for long-chain **dicarboxylic acid** prodn.)
- IT Genetic engineering  
(of *Candida tropicalis*, for long-chain **dicarboxylic acid** prodn.)
- IT **Carboxylic acids, preparation**

RL: PREP (Preparation)

(di-, long-chain, manuf. of, genetic engineering of Candida tropicalis  
for)

L17 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1985:61628 CAPLUS

DOCUMENT NUMBER: 102:61628

TITLE: Oxidative ether cleavage with p-nitroperbenzoic acid

AUTHOR(S): Schneider, Hans Joerg; Ahlhelm, Alfred; Mueller,  
Walter

CORPORATE SOURCE: Univ. Saarlandes, Saarbruecken, D-6600/11, Fed. Rep.  
Ger.

SOURCE: Chem. Ber. (1984), 117(11), 3297-302

CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal

LANGUAGE: German

AB The reaction of p-nitroperbenzoic acid in CHCl<sub>3</sub> with Me<sub>2</sub>CH(CH<sub>2</sub>)<sub>3</sub>OMe and (Me<sub>2</sub>CHCH<sub>2</sub>CH<sub>2</sub>)<sub>2</sub>O leads by selective attack at C-H bonds in the .  
alpha.-position to the ether oxygen to hemiacetals, which decomp.  
to aldehydes and alcs., yielding carboxylic acids.

Secondary alkoxy groups, as in ethoxycyclohexane, furnish Baeyer-Villiger oxidn. products of initially formed ketones. Kinetic measurements with substituted benzyl Me ethers show a Hammett reaction const. .rho. = -0.9, which is in accord with the obsd. relatively small discrimination between secondary and tertiary C-H bonds. The results are compared with similar hydroxylations of alkanes and with monooxygenase

reactions and point to oxenoid transition states. Radical reactions, as found with some alkanes are not obsd., which is shown by the small amts. of PhNO<sub>2</sub> (.1toreq.10%) formed during the reaction. <sup>13</sup>C-NMR shifts of several ethers and oxidn. products are reported.

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IT 123-51-3P 502-44-3P 503-74-2P 622-45-7P 626-89-1P 646-07-1P  
2412-73-9P 5299-60-5P 94368-15-7P 94368-16-8P

RL: PREP (Preparation)

(formation and carbon-13 NMR of)

L17 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1977:498893 CAPLUS

DOCUMENT NUMBER: 87:98893

TITLE: Biosynthesis of cutin. .omega.-

Hydroxylation of fatty acids by a microsomal preparation from germinating Vicia faba

Soliday, Charles L.; Kolattukudy, P. E.

AUTHOR(S): CORPORATE SOURCE: Dep. Agric. Chem., Washington State Univ., Pullman,  
Wash., USA

SOURCE: Plant Physiol. (1977), 59(6), 1116-21

CODEN: PLPHAY

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB .omega.-**Hydroxylation** of fatty acids, which is a key reaction in the biosynthesis of cutin and suberin, was demonstrated in a cell-free **prepn.** from a higher plant. A crude microsomal fraction (105,000-g pellet) from germinating embryonic shoots of *V. faba* catalyzed the conversion of palmitic acid to .omega.-hydroxypalmitic acid.

As the crude cell-free **prepn.** also catalyzes the formation of other hydroxy acids such as .alpha.- and .beta.-hydroxy acids, the .omega.-**hydroxylation** product was identified by gas chromatog. on a polyester column and reverse phase, high performance liq. chromatog., 2 techniques which were shown to resolve the positional isomers. Gas chromatog. anal. of the **dicarboxylic acid** obtained by CrO<sub>3</sub> oxidn. of the enzymic product also confirmed the identity

of the enzymic .omega.-**hydroxylation** product. This enzymic **hydroxylation** required O and NADPH, but substitution of NADH resulted in nearly half the reaction rate obtained with NADPH. Maximal rates of .omega.-**hydroxylation** occurred at pH 8 and the rate increased in a sigmoidal manner with increasing concns. of palmitic acid. This .omega.-**hydroxylation** was inhibited by the classical **mixed function oxidase** inhibitors such as metal chelators (o-phenanthroline, 8-hydroxyquinoline, and .alpha... .alpha.-dipyridyl), NaN<sub>3</sub>, and thiol reagents (N-ethylmaleimide and p-chloromercuribenzoate). As expected of a hydroxylase, involving cytochrome P450, the present o-hydroxylase was inhibited by CO and this enzyme system showed unusually high sensitivity to this inhibition; 10% CO caused inhibition and 30% CO completely inhibited the reaction. Another unusual feature was that the inhibition caused by any level of CO could not be reversed by light (420-460 nm).

TI Biosynthesis of cutin. .omega.-**Hydroxylation** of fatty acids by a microsomal **preparation** from germinating *Vicia faba*

AB .omega.-**Hydroxylation** of fatty acids, which is a key reaction in the biosynthesis of cutin and suberin, was demonstrated in a cell-free **prepn.** from a higher plant. A crude microsomal fraction (105,000-g pellet) from germinating embryonic shoots of *V. faba* catalyzed the conversion of palmitic acid to .omega.-hydroxypalmitic acid.

As the crude cell-free **prepn.** also catalyzes the formation of other hydroxy acids such as .alpha.- and .beta.-hydroxy acids, the .omega.-**hydroxylation** product was identified by gas chromatog. on a polyester column and reverse phase, high performance liq. chromatog., 2 techniques which were shown to resolve the positional isomers. Gas chromatog. anal. of the **dicarboxylic acid** obtained by CrO<sub>3</sub> oxidn. of the enzymic product also confirmed the identity

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high sensitivity to this inhibition; 10% CO caused inhibition and 30% CO completely inhibited the reaction. Another unusual feature was that the inhibition caused by any level of CO could not be reversed by light (420-460 nm).

ST **omega hydroxylation** fatty acid Vicia

IT Fatty acids, biological studies

RL: RCT (Reactant)

(.omega.-hydroxylation of, by microsome from broad  
bean)

IT Hydroxylation

(omega, of fatty acids by microsomal **prep.** from Vicia faba)

=> d full history

(FILE 'HOME' ENTERED AT 08:53:03 ON 10 MAY 2001)

FILE 'CAPLUS' ENTERED AT 08:53:11 ON 10 MAY 2001  
E .BETA.-OXIDN.-BLOCKED/CT  
E E2  
E E3+ALL  
E .BETA.-OXIDATION/CT  
E E3+ALL

FILE 'REGISTRY' ENTERED AT 09:37:48 ON 10 MAY 2001

L1 1 SEA ABB=ON PLU=ON 9038-14-6/RN  
D  
L2 1 SEA ABB=ON PLU=ON 9023-03-4/RN  
D

FILE 'CAOLD, CAPLUS, CROPU, DGENE, DPCI, ENCOMPPAT, ENCOMPPAT2,  
EUROPATFULL, IFIPAT, INPADOC, JAPIO, PAPERCHEM2, PATDD, PATDPA, PATOSDE,  
PATOSEP, PATOSWO, PCTFULL, PIRA, RAPRA, SYNTHLINE, TULSA, TULSA2,  
USPATFULL, WPIDS' ENTERED AT 09:39:31 ON 10 MAY 2001

FILE 'REGISTRY' ENTERED AT 09:39:57 ON 10 MAY 2001

L3 SET SMARTSELECT ON  
SEL PLU=ON L1 1- CHEM : 26 TERMS  
SET SMARTSELECT OFF

FILE 'CAOLD, CAPLUS, CROPU, DGENE, DPCI, ENCOMPPAT, ENCOMPPAT2,  
EUROPATFULL, IFIPAT, INPADOC, JAPIO, PAPERCHEM2, PATDD, PATDPA, PATOSDE,  
PATOSEP, PATOSWO, PCTFULL, PIRA, RAPRA, SYNTHLINE, TULSA, TULSA2,  
USPATFULL, WPIDS' ENTERED AT 09:40:22 ON 10 MAY 2001

FILE 'REGISTRY' ENTERED AT 09:40:25 ON 10 MAY 2001

L4 SET SMARTSELECT ON  
SEL PLU=ON L2 1- CHEM : 16 TERMS  
SET SMARTSELECT OFF

FILE 'CAOLD, CAPLUS, CROPU, DGENE, DPCI, ENCOMPPAT, ENCOMPPAT2,  
EUROPATFULL, IFIPAT, INPADOC, JAPIO, PAPERCHEM2, PATDD, PATDPA, PATOSDE,  
PATOSEP, PATOSWO, PCTFULL, PIRA, RAPRA, SYNTHLINE, TULSA, TULSA2,  
USPATFULL, WPIDS' ENTERED AT 09:40:33 ON 10 MAY 2001

L5 18788 SEA ABB=ON PLU=ON L3  
L6 7868 SEA ABB=ON PLU=ON L4  
L7 25412 SEA ABB=ON PLU=ON L5 OR L6  
L8 830900 SEA ABB=ON PLU=ON DICARBOXYLIC ACID# OR (CARBOXYLIC ACIDS  
(L) DICARBOXYLIC) OR CARBOXYLIC ACID# OR MONOCARBOXYLIC ACID#  
L9 573 SEA ABB=ON PLU=ON L7 AND L8  
L10 361 SEA ABB=ON PLU=ON L9 AND (PREPAR? OR SYNTHE? OR MAK? OR  
PREP/RL)  
L11 11119 SEA ABB=ON PLU=ON (HYDROXYLATION (L) .OMEGA.-HYDROXYLATION)  
OR (HYDROXYLATION (L) ALKANE#) OR (.ALPHA.-HYDROXYLATION) OR  
(HYDROXYLATION (L) .ALPHA.)  
L12 76 SEA ABB=ON PLU=ON L11 AND L10  
L13 74 DUP REM L12 (2 DUPLICATES REMOVED)  
L14 1066 SEA ABB=ON PLU=ON CANDIDA MALTOSA OR CANDIDA CLOACAE OR  
CANDIDA NOVELLUS OR CANDIDA SUBTROPICALIS  
L15 7 SEA ABB=ON PLU=ON L14 AND L10  
L16 6 DUP REM L15 (1 DUPLICATE REMOVED)  
D IBIB AB 1-6

D IBIB AB 1  
D IBIB AB 1-6

=> d ibib ab 1-6

L16 ANSWER 1 OF 6 PCTFULL COPYRIGHT 2001 MicroPatent  
ACCESSION NUMBER: 2000065061 PCTFULL EW 200044 ED 20001124  
TITLE (ENGLISH): NUCLEIC ACID SEQUENCES FROM <i> CANDIDA </i> YEASTS  
WHICH CODE CYTOCHROME B5 POLYPEPTIDES  
TITLE (FRENCH): SEQUENCES D'ACIDE NUCLEIQUE ISSUES DE LEVURES DE <i>  
CANDIDA </i>, CODANT DES POLYPEPTIDES B5 CYTOCHROMES  
TITLE (GERMAN): NUKLEINSAEURE-SEQUENZEN AUS <i> CANDIDA </i> HEFEN,  
DIE CYTOCHROM B5-POLYPEPTIDE KODIEREN  
INVENTOR(S): SCHUNCK, Wolf-Hagen; CHERNOGOLOV, Alexei  
PATENT ASSIGNEE(S): MAX-DELBURUECK-CENTRUM FUeR MOLEKULARE MEDIZIN  
LANGUAGE OF PUBL.: German  
LANGUAGE OF FILING: German  
DOCUMENT TYPE: Patent  
PATENT INFORMATION:

	NUMBER	KIND	DATE
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DESIGNATED STATES: WO 2000065061 A2 20001102  
AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ  
DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP  
KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW  
MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ  
UA UG US UZ VN YU ZA ZW GH GM KE LS MW SD SL SZ TZ UG  
ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI  
FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN  
GW ML MR NE SN TD TG

APPLICATION INFO.: WO 2000-DE1246 20000418

PRIORITY (ORIGINAL): DE 1999-199 18 763.0 19990424

ABEN The invention relates to nucleic acid sequences from <i> Candida </i> yeasts, preferably from <i> Candida maltosa </i>, which code cytochrome b5 polypeptides, as well as to the corresponding cytochrome b5 polypeptides and their use for increasing the activity of cytochrome P450 systems, especially for stimulating the activity of alkane-hydroxylating and fatty acid-hydroxylating cytochrome P450 systems during the production of long-chained dicarboxylic acids (#ge#C10).

ABFR L'invention concerne des sequences d'acide nucleique issues de levures de <i> Candida </i>, de preference de <i> Candida maltosa </i>, qui codent des polypeptides b5 cytochromes. L'invention concerne egalement les polypeptides b5 cytochromes correspondants et leur utilisation pour augmenter l'activite de systemes P450 cytochromes, notamment pour stimuler l'activite de systemes P450 cytochromes a effet hydroxylant de l'alcanes et de l'acide gras lors de la preparation d'acides dicarboxyliques a chaine longue (#ge#C10).

ABDE Die Erfindung betrifft Nukleinsaeure-Sequenzen aus <i> Candida </i> Hefen vorzugsweise aus <i> Candida maltosa </i>, die Cytochrom b5-Polypeptide kodieren sowie die entsprechenden Cytochrom b5-Polypeptide und ihre Verwendung zur Erhoehung der Aktivitaet von Cytochrom P450 Systemen, insbesondere zur Stimulierung der Aktivitaet Alkan- und Fettsaeure-hydroxylierender Cytochrom P450 Systeme bei der Produktion langkettiger Dicarbonsaeuren (#ge#C10).

L16 ANSWER 2 OF 6 PCTFULL COPYRIGHT 2001 MicroPatent  
 ACCESSION NUMBER: 2000034473 PCTFULL EW 200024 ED 20000712  
 TITLE (ENGLISH): SEVEN TRANSMEMBRANE DOMAIN RECEPTOR ZSIG56  
 TITLE (FRENCH): DOMAINE TRANSMEMBRANAIRE 7 ZSIG56  
 INVENTOR(S): SHEPPARD, Paul, O.; ELLSWORTH, Jeff, L.  
 PATENT ASSIGNEE(S): ZYMOGENETICS, INC.  
 LANGUAGE OF PUBL.: English  
 LANGUAGE OF FILING: English  
 DOCUMENT TYPE: Patent  
 PATENT INFORMATION:  
 DESIGNATED STATES:  
 NUMBER KIND DATE  
 -----
 WO 2000034473 A2 20000615  
 AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE  
 DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE  
 KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO  
 NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG UZ  
 VN YU ZA ZW GH GM KE LS MW SD SL SZ TZ UG ZW AM AZ BY  
 KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE  
 IT LU MC NL PT SE BF BJ CF CG CI CM GA GN GW ML MR NE  
 SN TD TG  
 APPLICATION INFO.: WO 1999-US28492 19991202  
 PRIORITY (ORIGINAL): US 1998-09/208691 19981210  
 ABEN The present invention relates to polynucleotide and polypeptide molecules for a seven transmembrane domain receptor designated zsig56. The polypeptides, and polynucleotides encoding them are useful treating pathological conditions in such diverse tissue, as kidney, thyroid, gastrointestinal, CSN and reproductive. Such conditions include hypertension, hyper- and hypothyroidism, neurotransmission, gastrointestinal motility, inflammation and reproduction. The present invention also includes antibodies to the zsig56 polypeptides.  
 ABFR L'invention concerne des molecules polynucleotidiques et polypeptidiques pour un recepteur du domaine transmembranaire 7 appele zsig56. Les polypeptides et les polynucleotides codant pour eux conviennent pour le traitement d'etats pathologiques de tissus aussi divers que les reins, la glande thyroide, le tube digestif, le systeme nerveux central et l'appareil genital. Lesdits etats pathologiques concernent l'hypertension, l'hyperthyroïdie, l'hypothyroïdie, la neurotransmission, le transit intestinal, l'inflammation et la reproduction. L'invention concerne également des anticorps dirigés contre les polypeptides zsig56.  
 L16 ANSWER 3 OF 6 PCTFULL COPYRIGHT 2001 MicroPatent  
 ACCESSION NUMBER: 2000020566 PCTFULL EW 200015 ED 20000503  
 TITLE (ENGLISH): CYTOCHROME P450 MONOOXYGENASE AND NADPH  
 CYTOCHROME P450  
 OXIDOREDUCTASE GENES AND PROTEINS RELATED TO THE  
 OMEGA  
 HYDROXYLASE  
 COMPLEX OF *CANDIDA TROPICALIS* AND METHODS  
 RELATING THERETO  
 TITLE (FRENCH): GENES DE LA MONOOXYGENASE DU CYTOCHROME P450  
 ET DE  
 L'OXYDOREDUCTASE NADPH DU CYTOCHROME P450 ET  
 PROTEINES  
 ASSOCIEES AU  
 COMPLEXE DE L'OMEGA HYDROXYLASE DE *CANDIDA TROPICALIS* ET PROCEDES  
 ASSOCIES  
 INVENTOR(S): WILSON, C., Ron; CRAFT, David, L.; EIRICH, L.,  
 Dudley;

ESHOO, Mark; MADDURI, Krishna, M.; CORNETT, Cathy,  
A.;  
BRENNER, Alfred, A.; TANG, Maria; LOPER, John, C.;  
GLEESON, Martin  
PATENT ASSIGNEE(S): HENKEL CORPORATION  
LANGUAGE OF PUBL.: English  
LANGUAGE OF FILING: English  
DOCUMENT TYPE: Patent  
PATENT INFORMATION:

	NUMBER	KIND	DATE
DESIGNATED STATES:	WO 2000020566	A2	20000413
	AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW GH GM KE LS MW SD SL SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN GW ML MR NE SN TD TG		
APPLICATION INFO.:	WO 1999-US20797		19990910
PRIORITY (ORIGINAL):	US 1998-60/103099		19981005
	US 1999-		19990310

ABEN Novel genes have been isolated which encode cytochrome P450 and NADPH reductase enzyme of the #ohgr#-hydroxylase complex of <i>C. tropicalis</i> 20336. Vectors including these genes, transfected host cells and transformed host cells are provided. Methods of producing of cytochrome P450 and NADPH reductase enzymes are also provided which involve transforming a host cell with a gene encoding these enzymes and culturing the cells. Methods of increasing the production of a **dicarboxylic acid** and methods of increasing production of the aforementioned enzymes are also provided which involve increasing in the host cell the number of genes encoding these enzymes. A method for discriminating members of a gene family by quantifying the expression of genes is also provided.

ABFR La presente invention concerne des genes isolés qui codent pour des enzymes du cytochrome P450 et de reductase de NADPH du complexe #ohgr#-hydroxylase de <i>C. tropicalis</i> 20336. L'invention concerne aussi des vecteurs contenant ces genes, des cellules hôtes transfectées ainsi que des cellules hôtes transformées. Sont aussi décrits des procédés de production des enzymes du cytochrome P450 et de reductase de NADPH qui consistent à transformer une cellule hôte à l'aide d'un gène qui code pour ces enzymes et à cultiver les cellules. L'invention concerne aussi des procédés d'augmentation de la production d'un acide dicarboxylique, des procédés d'augmentation de la production des enzymes mentionnées ci-dessus qui consistent à augmenter le nombre de gènes qui codent pour ces enzymes dans les cellules hôtes, ainsi qu'une méthode destinée à distinguer des éléments d'une famille de gènes en quantifiant l'expression des gènes.

L16 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2001 ACS DUPLICATE 1  
ACCESSION NUMBER: 1999:77701 CAPLUS  
DOCUMENT NUMBER: 130:138397  
TITLE: Yeasts with elevated cytochrome P450 levels and their use in the manufacture of monotermal and diterminal aliphatic carboxylates from alkanes  
INVENTOR(S): Fallon; Robert D.; Payne, Mark S.; Picataggio, Stephen K.; Wu, Shijun

PATENT ASSIGNEE(S): E.I. Du Pont De Nemours and Company, USA  
 SOURCE: PCT Int. Appl., 58 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9904014	A2	19990128	WO 1998-US14935	19980720
WO 9904014	A3	19990520		
W: AU, CA, IL, JP, NZ, US				
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
AU 9884982	A1	19990210	AU 1998-84982	19980720
EP 1003881	A2	20000531	EP 1998-935806	19980720
R: DE, DK, ES, FR, GB, IT, NL, SE, IE				
PRIORITY APPLN. INFO.:			US 1997-53215	P 19970721
			WO 1998-US14935	W 19980720

AB Strains of *Pichia pastoris* and *Candida maltosa* that have increased cytochrome P 450 activity or lack the .beta.-oxidn. pathway that can be used to convert c6-C22 alkanes to monoterinal and diterinal carboxylic acids are described. Expression constructs carrying cytochrome P 450 genes under the control of powerful inducible promoters (AOX1, PGK) were constructed by std. methods and introduced into *P. pastoris* or *C. maltosa* to give novel or increased P 450 activities. Disruption of .beta.-oxidn. in *C. maltosa* was achieved by insertional inactivation of the POX4 gene. A strain of *C. maltosa* cultured in a complete medium that was supplemented with dodecane 20 g/L when glucose was almost completely depleted was able to generate dodecanedioic acid at 3.4 g/h for 51 h.

L16 ANSWER 5 OF 6	PCTFULL	COPYRIGHT 2001 MicroPatent	
ACCESSION NUMBER:	1998055612	PCTFULL	
TITLE (ENGLISH):	NEUROKININ B PRECURSORS		
TITLE (FRENCH):	PRECURSEURS DE NEUROKININE B		
INVENTOR(S):	SHEPPARD, Paul, O.		
PATENT ASSIGNEE(S):	ZYMOGENETICS, INC.		
LANGUAGE OF PUBL.: English			
LANGUAGE OF FILING: English			
DOCUMENT TYPE: Patent			
PATENT INFORMATION:	NUMBER	KIND	DATE
	WO 9855612	A1	19981210
DESIGNATED STATES:	AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG UZ VN ZW GH GM KE LS MW SD SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN ML MR NE SN TD TG		
APPLICATION INFO.:	WO 1998-US10842		19980528
PRIORITY (ORIGINAL):	US 1997-60/048290		19970602
ABEN	The present invention relates to zneurok1 polypeptides and polynucleotides encoding the same. These polypeptides are novel members of a family of proteins that are precursors of neurokinin B, a ten amino		

acid moiety of biological significance. The polypeptides, and polynucleotides encoding them, are useful in the study of prohormone convertase function and neurokinin receptors. The present invention also includes antibodies to the zneurok1 polypeptides.

ABFR Cette invention concerne des polypeptides zneurok1 ainsi que des polynucleotides codant ces derniers. Ces polypeptides consistent en de nouveaux membres d'une famille de proteines qui sont des precurseurs de neurokinine B, un fragment de l'acide amine dix jouant un role biologique important. Ces polypeptides, ainsi que les polynucleotides qui les codent, sont utiles lors de l'etude de la fonction de convertase de prohormone, ainsi que des recepteurs de neurokinine. Cette invention concerne egalement des anticorps diriges contre les polypeptides zneurok1.

L16 ANSWER 6 OF 6 PCTFULL COPYRIGHT 2001 MicroPatent  
ACCESSION NUMBER: 1996027678 PCTFULL  
TITLE (ENGLISH): PROCESS FOR HYDROXYLATING LONG-CHAIN ALKANES, FATTY ACIDS AND OTHER ALKYL COMPOUNDS  
TITLE (FRENCH): PROCEDE D'HYDROXYLATION D'ALCANES A LONGUE CHAINE, D'ACIDES GRAS ET D'AUTRES COMPOSES ALKYLES  
INVENTOR(S): ZIMMER, Thomas; KAMINSKI, Kristina; SCHUNCK, Wolf-Hagen; KAeRGEL, Eva; SCHELLER, Ulrich; MAUERSBERGER, Stephan  
PATENT ASSIGNEE(S): MAX-DELBRUECK-CENTRUM FUeR MOLEKULARE MEDIZIN; ZIMMER, Thomas; KAMINSKI, Kristina; SCHUNCK, Wolf-Hagen; KAeRGEL, Eva; SCHELLER, Ulrich; MAUERSBERGER, Stephan  
LANGUAGE OF PUBL.: German  
DOCUMENT TYPE: Patent  
PATENT INFORMATION:

	NUMBER	KIND	DATE
DESIGNATED STATES:	WO 9627678	A1	19960912
	JP US AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE		
APPLICATION INFO.:	WO 1996-DE410		19960301
PRIORITY (ORIGINAL):	DE 1995-195 07 546.3		19950303
ABEN	A microbial hydroxylation process is disclosed for selectively oxidising regions of long-chain alkanes, fatty acids and other alkyl compounds. The process should be easy to carry out and produce good yields of oxidation products, in particular hydroxylated fatty acids and long-chain <b>dicarboxylic acids</b> . The object of the invention is to modify yeast by genetic engineering so that when it is cultivated it expresses the required enzymes. The disclosed process is characterised in that the long-chain alkanes, fatty acids and other alkyl compounds are treated with <b>monooxygenase</b> systems that consist of cytochrome P450 and NADPH cytochrome P450 reductase, and the hydroxylation products are then separated. The <b>monooxygenase</b> systems are produced in the reaction mixture by simultaneous expression of their components in yeast, preferably <i>saccharomyces cerevisiae</i> . The invention relates essentially to a vector for modifying <i>saccharomyces</i> by genetic engineering. On the basis of the structure Yep 51, the vector contains reductase cDNA between the restriction sites SalK and BamHI and a second expression cassette bound in the restriction site NruI. The expression cassette consists of the GAL10 promoter, the sequence that codes for cytochrome		

P450 and the ADH1 terminator.

ABF L'invention concerne un procede d'hydroxylation microbienne qui sert a oxyder selectivement des regions d'alcanes a longue chaine, d'acides gras et d'autres composes alkyle. Le procede doit permettre d'obtenir de maniere aisee un bon rendement en produits d'oxydation, notamment des acides gras hydroxyles et des acides dicarboxyliques a longue chaine. L'invention a pour objet de modifier par genie genetique des levures de sorte qu'elles expriment les enzymes requises lorsqu'elles sont cultivees. Le procede se caracterise en ce que l'on traite les alcanes a longue chaine, les acides gras et les autres composes alkyle avec des systemes a **monooxygenase** constitues de cytochrome P450 et de NADPH-cytochrome P450-reductase, les produits d'hydroxylation etant ensuite isolés. Les systemes a **monooxygenase** sont produits dans le melange de reaction par expression simultanee de leurs composants dans des levures, de preference la saccharomyces cerevisiae. L'invention porte essentiellement sur un vecteur de modification par genie genetique de saccharomyces, qui sur la base de la structure fondamentale Yep 51 contient l'ADNc de la reductase entre les sites de restriction SalI et BamHI et une deuxième cassette d'expression liee dans le site de restriction NruI et constituee du promoteur GAL10, de la sequence de codage de cytochrome P450 et du terminateur ADH1.

=> d full history

(FILE 'HOME' ENTERED AT 11:17:32 ON 10 MAY 2001)

FILE 'REGISTRY' ENTERED AT 11:17:52 ON 10 MAY 2001

L1 1 SEA ABB=ON PLU=ON 9038-14-6/RN

D

L2 1 SEA ABB=ON PLU=ON 9023-03-4/RN

D

FILE 'HCAPLUS' ENTERED AT 11:18:36 ON 10 MAY 2001

FILE 'REGISTRY' ENTERED AT 11:18:51 ON 10 MAY 2001

SET SMARTSELECT ON

L3 SEL PLU=ON L1 1- CHEM : 26 TERMS  
SET SMARTSELECT OFF

FILE 'HCAPLUS' ENTERED AT 11:18:52 ON 10 MAY 2001

FILE 'REGISTRY' ENTERED AT 11:18:53 ON 10 MAY 2001

SET SMARTSELECT ON

L4 SEL PLU=ON L2 1- CHEM : 16 TERMS  
SET SMARTSELECT OFF

FILE 'HCAPLUS' ENTERED AT 11:18:54 ON 10 MAY 2001

L5 15271 SEA ABB=ON PLU=ON L3

L6 7478 SEA ABB=ON PLU=ON L4

L7 21584 SEA ABB=ON PLU=ON L5 OR L6

L8 188643 SEA ABB=ON PLU=ON DICARBOXYLIC ACID# OR (CARBOXYLIC ACIDS  
(L) DICARBOXYLIC) OR CARBOXYLIC ACID# OR MONOCARBOXYLIC ACID#

L9 5883 SEA ABB=ON PLU=ON (HYDROXYLATION (L) .OMEGA.-HYDROXYLATION)  
OR (HYDROXYLATION (L) ALKANE#) OR (.ALPHA.-HYDROXYLATION) OR  
(HYDROXYLATION (L) .ALPHA.)

L10 1338 SEA ABB=ON PLU=ON PICHIA PASTORIS

L11 4 SEA ABB=ON PLU=ON L7 (L) L8 (L) L9

D IBIB AB 1-4

D IBIB AB HIT 1-4

L12 0 SEA ABB=ON PLU=ON L11 AND L10

=> d ibib ab 1-4

L11 ANSWER 1 OF 4 HCPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1993:5685 HCPLUS  
DOCUMENT NUMBER: 118:5685  
TITLE: Metabolic engineering of *Candida tropicalis* for the production of long-chain dicarboxylic acids  
AUTHOR(S): Picataggio, Stephen; Rohrer, Tracy; Deanda, Kristine; Lanning, Dawn; Reynolds, Robert; Mielenz, Jonathan; Eirich, L. Dudley  
CORPORATE SOURCE: Microb. Technol. Dep., Cognis Inc., Santa Rosa, CA, 95407, USA  
SOURCE: Bio/Technology (1992), 10(8), 894-8  
CODEN: BTCHDA; ISSN: 0733-222X  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB An industrial strain of the yeast *C. tropicalis* was engineered for the efficient prodn. of long-chain **dicarboxylic acids**, which are important raw materials for the chem. industry. By sequential disruption of the 4 genes encoding both isoenzymes of the acyl-CoA oxidase which catalyzes the first reaction in the .beta.-oxidn. pathway, **alkane** and fatty acid substrates were successfully redirected to the .omega.-oxidn. pathway. Consequently, the conversion efficiency and chem. selectivity of their terminal oxidn. to the corresponding **dicarboxylic acids** was improved to 100%. The specific productivity of the bioconversion was increased further by amplification of the **cytochrome P 450 monooxygenase** and NADPH-cytochrome reductase genes encoding the rate-limiting .omega.-hydroxylase in the .omega.-oxidn. pathway. The amplified strains demonstrated increased .omega.-hydroxylase activity and a 30% increase in productivity compared to the .beta.-oxidn.-blocked strain in fermns. The bioconversion is effective for the selective terminal oxidn. of both satd. and unsatd. linear aliph. substrates with C12-22 chain-lengths and also avoids the undesirable chain modifications assocd. with passage through the .beta.-oxidn. pathway, such as unsatn., **hydroxylation**, or chain shortening. It is now possible to efficiently produce a wide range of previously unavailable satd. and unsatd. **dicarboxylic acids** with a high degree of purity.

L11 ANSWER 2 OF 4 HCPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1991:220705 HCPLUS  
DOCUMENT NUMBER: 114:220705  
TITLE: Microsomal oxidation of dodecylthioacetic acid (a 3-thia fatty acid) in rat liver  
AUTHOR(S): Hvattum, Erlend; Bergseth, Steinar; Pedersen, Catharina N.; Bremer, Jon; Aarsland, Asle; Berge, Rolf K.  
CORPORATE SOURCE: Inst. Med. Biochem., Univ. Oslo, Oslo, 0317, Norway  
SOURCE: Biochem. Pharmacol. (1991), 41(6-7), 945-53  
CODEN: BCPCA6; ISSN: 0006-2952  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB [1-14C]Dodecylthioacetic acid (DTA), a 3-thia fatty acid, is .omega. (.omega.-1)-hydroxylated and sulfur oxygenated at about equal rates in rat liver microsomes. In prolonged incubations DTA is converted to

.omega.-hydroxydodecylsulfoxyacetic acid. .omega.-  
**Hydroxylation** of DTA is catalyzed by cytochrome P450IVA1 (or a very closely related isoenzyme in the same gene family), the fatty acid .omega.-hydroxylating enzyme. It is absolutely dependent on NADPH and inhibited by CO, and lauric acid is a competing substrate. .omega.  
.Hydroxylation of DTA is increased by feeding tetradecylthioacetic acid (TTA), a 3-thia fatty acid, for 4 days to rats. .omega.-**Hydroxylation** of [1-14C] lauric acid is also induced by TTA and other 3-thia **carboxylic acids**. A close relationship was obsd. between induction of microsomal .omega.-**hydroxylation** of fatty acid and palmitoyl-CoA hydrolase activity. DTA is .omega.-hydroxylated at about the same rate

as

the physiol. substrate lauric acid. The sulfur oxygenation of DTA is catalyzed by liver microsomal **flavin-contg.** **monooxygenase** (FMO) (EC 1.14.13.8). It is dependent on either NADH or 1NADPH. The Km value for NADH was .apprx.five times larger than the Km value for NADPH. It is inhibited by methimazole and not affected by CO. It is not induced by TTA.

L11 ANSWER 3 OF 4 HCPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1985:61628 HCPLUS

DOCUMENT NUMBER: 102:61628

TITLE: Oxidative ether cleavage with p-nitroperbenzoic acid Schneider, Hans Joerg; Ahlhelm, Alfred; Mueller, Walter

AUTHOR(S): Univ. Saarlandes, Saarbruecken, D-6600/11, Fed. Rep. Ger.

CORPORATE SOURCE: Chem. Ber. (1984), 117(11), 3297-302  
CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal

LANGUAGE: German

AB The reaction of p-nitroperbenzoic acid in CHCl<sub>3</sub> with Me<sub>2</sub>CH(CH<sub>2</sub>)<sub>3</sub>OMe and (Me<sub>2</sub>CHCH<sub>2</sub>CH<sub>2</sub>)<sub>2</sub>O leads by selective attack at C-H bonds in the .alpha.-position to the ether oxygen to hemiacetals, which decomp. to aldehydes and alcs., yielding **carboxylic acids**. Secondary alkoxy groups, as in ethoxycyclohexane, furnish Baeyer-Villiger oxidn. products of initially formed ketones. Kinetic measurements with substituted benzyl Me ethers show a Hammett reaction const. .rho. = -0.9, which is in accord with the obsd. relatively small discrimination between secondary and tertiary C-H bonds. The results are compared with similar **hydroxylations** of alkanes and with **monooxygenase** reactions and point to oxenoid transition states. Radical reactions, as found with some **alkanes** are not obsd., which is shown by the small amts. of PhNO<sub>2</sub> (.1toreq.10%) formed during the reaction. <sup>13</sup>C-NMR shifts of several ethers and oxidn. products are reported.

L11 ANSWER 4 OF 4 HCPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1977:498893 HCPLUS

DOCUMENT NUMBER: 87:98893

TITLE: Biosynthesis of cutin. .omega.-**Hydroxylation** of fatty acids by a microsomal preparation from germinating

AUTHOR(S): Vicia faba Soliday, Charles L.; Kolattukudy, P. E.

CORPORATE SOURCE: Dep. Agric. Chem., Washington State Univ., Pullman, Wash., USA

SOURCE: Plant Physiol. (1977), 59(6), 1116-21  
CODEN: PLPHAY

DOCUMENT TYPE: Journal

LANGUAGE: English

AB .omega.-Hydroxylation of fatty acids, which is a key reaction in the biosynthesis of cutin and suberin, was demonstrated in a cell-free prepn. from a higher plant. A crude microsomal fraction (105,000-g pellet) from germinating embryonic shoots of V. faba catalyzed the conversion of palmitic acid to .omega.-hydroxypalmitic acid. As the crude cell-free prepn. also catalyzes the formation of other hydroxy acids such as .alpha.- and .beta.-hydroxy acids, the .omega.-hydroxylation product was identified by gas chromatog. on a polyester column and reverse phase, high performance liq. chromatog., 2 techniques which were shown to resolve the positional isomers. Gas chromatog. anal. of the dicarboxylic acid obtained by CrO<sub>3</sub> oxidn. of the enzymic product also confirmed the identity of the enzymic .omega.-hydroxylation product. This enzymic hydroxylation required O<sub>2</sub> and NADPH, but substitution of NADH resulted in nearly half the reaction rate obtained with NADPH. Maximal rates of .omega.-hydroxylation occurred at pH 8 and the rate increased in a sigmoidal manner with increasing concns. of palmitic acid. This .omega.-hydroxylation was inhibited by the classical mixed function oxidase inhibitors such as metal chelators (o-phenanthroline, 8-hydroxyquinoline, and .alpha.,.alpha.-dipyridyl), NaN<sub>3</sub>, and thiol reagents (N-ethylmaleimide and p-chloromercuribenzoate). As expected of a hydroxylase, involving cytochrome P450, the present o-hydroxylase was inhibited by CO and this enzyme system showed unusually high sensitivity to this inhibition; 10% CO caused inhibition and 30% CO completely inhibited the reaction. Another unusual feature was that the inhibition caused by any level of CO could not be reversed by light (420-460 nm).

=> d ibib ab hit 1-4

L11 ANSWER 1 OF 4 HCPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1993:5685 HCPLUS  
DOCUMENT NUMBER: 118:5685  
TITLE: Metabolic engineering of *Candida tropicalis* for the production of long-chain dicarboxylic acids  
AUTHOR(S): Picataggio, Stephen; Rohrer, Tracy; Deanda, Kristine; Lanning, Dawn; Reynolds, Robert; Mielenz, Jonathan; Eirich, L. Dudley  
CORPORATE SOURCE: Microb. Technol. Dep., Cognis Inc., Santa Rosa, CA, 95407, USA  
SOURCE: Bio/Technology (1992), 10(8), 894-8  
CODEN: BTCHDA; ISSN: 0733-222X  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB An industrial strain of the yeast *C. tropicalis* was engineered for the efficient prodn. of long-chain dicarboxylic acids, which are important raw materials for the chem. industry. By sequential disruption of the 4 genes encoding both isoenzymes of the acyl-CoA oxidase which catalyzes the first reaction in the .beta.-oxidn. pathway, alkane and fatty acid substrates were successfully redirected to the .omega.-oxidn. pathway. Consequently, the conversion efficiency and chem. selectivity of their terminal oxidn. to the corresponding dicarboxylic acids was improved to 100%. The specific productivity of the bioconversion was increased further by amplification of the cytochrome P 450

**monooxygenase** and NADPH-cytochrome reductase genes encoding the rate-limiting .omega.-hydroxylase in the .omega.-oxidn. pathway. The amplified strains demonstrated increased .omega.-hydroxylase activity and a 30% increase in productivity compared to the .beta.-oxidn.-blocked strain in fermns. The bioconversion is effective for the selective terminal oxidn. of both satd. and unsatd. linear aliph. substrates with C12-22 chain-lengths and also avoids the undesirable chain modifications assocd. with passage through the .beta.-oxidn. pathway, such as unsatn., **hydroxylation**, or chain shortening. It is now possible to efficiently produce a wide range of previously unavailable satd. and unsatd. **dicarboxylic acids** with a high degree of purity.

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L11 ANSWER 2 OF 4 HCPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1991:220705 HCPLUS

DOCUMENT NUMBER: 114:220705

TITLE: Microsomal oxidation of dodecylthioacetic acid (a 3-thia fatty acid) in rat liver

AUTHOR(S): Hvattum, Erlend; Bergseth, Steinar; Pedersen, Catharina N.; Bremer, Jon; Aarsland, Asle; Berge, Rolf

K.

CORPORATE SOURCE: Inst. Med. Biochem., Univ. Oslo, Oslo, 0317, Norway

SOURCE: Biochem. Pharmacol. (1991), 41(6-7), 945-53

CODEN: BCPCA6; ISSN: 0006-2952

DOCUMENT TYPE: Journal

LANGUAGE: English

AB [1-14C]Dodecylthioacetic acid (DTA), a 3-thia fatty acid, is .omega. (.omega.-1)-hydroxylated and sulfur oxygenated at about equal rates in rat

liver microsomes. In prolonged incubations DTA is converted to .omega.-hydroxydodecylsulfoxyacetic acid. .omega.-**Hydroxylation** of DTA is catalyzed by cytochrome P450IVA1 (or a very closely related isoenzyme in the same gene family), the fatty acid .omega.-hydroxylating enzyme. It is absolutely dependent on NADPH and inhibited by CO, and lauric acid is a competing substrate. .omega.

.-Hydroxylation of DTA is increased by feeding tetradecylthioacetic acid (TTA), a 3-thia fatty acid, for 4 days to rats. .omega.-Hydroxylation of [1-14C] lauric acid is also induced by TTA and other 3-thia carboxylic acids. A close relationship was obsd. between induction of microsomal .omega.-hydroxylation of fatty acid and palmitoyl-CoA hydrolase activity. DTA is .omega.-hydroxylated at about the same rate

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the physiol. substrate lauric acid. The sulfur oxygenation of DTA is catalyzed by liver microsomal flavin-contg. monooxygenase (FMO) (EC 1.14.13.8). It is dependent on either NADH or 1NADPH. The Km value for NADH was .apprx. five times larger than the Km value for NADPH. It is inhibited by methimazole and not affected by CO. It is not induced by TTA.

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L11 ANSWER 3 OF 4 HCPLUS COPYRIGHT 2001 ACS  
ACCESSION NUMBER: 1985:61628 HCPLUS  
DOCUMENT NUMBER: 102:61628  
TITLE: Oxidative ether cleavage with p-nitroperbenzoic acid  
AUTHOR(S): Schneider, Hans Joerg; Ahlhelm, Alfred; Mueller, Walter  
CORPORATE SOURCE: Univ. Saarlandes, Saarbruecken, D-6600/11, Fed. Rep. Ger.  
SOURCE: Chem. Ber. (1984), 117(11), 3297-302  
CODEN: CHBEAM; ISSN: 0009-2940  
DOCUMENT TYPE: Journal  
LANGUAGE: German  
AB The reaction of p-nitroperbenzoic acid in CHCl<sub>3</sub> with Me<sub>2</sub>CH(CH<sub>2</sub>)<sub>3</sub>OMe and (Me<sub>2</sub>CHCH<sub>2</sub>CH<sub>2</sub>)<sub>2</sub>O leads by selective attack at C-H bonds in the .alpha.-position to the ether oxygen to hemiacetals, which decomp. to aldehydes and alcs., yielding carboxylic acids. Secondary alkoxy groups, as in ethoxycyclohexane, furnish Baeyer-Villiger oxidn. products of initially formed ketones. Kinetic measurements with substituted benzyl Me ethers show a Hammett reaction const. .rho. = -0.9, which is in accord with the obsd. relatively small discrimination between secondary and tertiary C-H bonds. The results are compared with similar hydroxylations of alkanes and with monooxygenase

reactions and point to oxenoid transition states. Radical reactions, as found with some **alkanes** are not obsd., which is shown by the small amts. of PhNO<sub>2</sub> (.1toreq.10%) formed during the reaction. <sup>13</sup>C-NMR shifts of several ethers and oxidn. products are reported.

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L11 ANSWER 4 OF 4 HCPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1977:498893 HCPLUS

DOCUMENT NUMBER: 87:98893

TITLE: Biosynthesis of cutin. **.omega.-Hydroxylation of fatty**

acids by a microsomal preparation from germinating Vicia faba

AUTHOR(S): Soliday, Charles L.; Kolattukudy, P. E.

CORPORATE SOURCE: Dep. Agric. Chem., Washington State Univ., Pullman, Wash., USA

SOURCE: Plant Physiol. (1977), 59(6), 1116-21

CODEN: PLPHAY

DOCUMENT TYPE: Journal

LANGUAGE: English

AB **.omega.-Hydroxylation** of fatty acids, which is a key reaction in the biosynthesis of cutin and suberin, was demonstrated in a cell-free prepn. from a higher plant. A crude microsomal fraction (105,000-g pellet) from germinating embryonic shoots of *V. faba* catalyzed the conversion of palmitic acid to **.omega.-hydroxypalmitic acid**. As the crude cell-free prepn. also catalyzes the formation of other hydroxy acids

such as **.alpha.-** and **.beta.-hydroxy acids**, the **.omega.-hydroxylation** product was identified by gas chromatog. on a polyester column and reverse phase, high performance liq. chromatog., 2 techniques which were shown to resolve the positional isomers. Gas chromatog. anal. of the **dicarboxylic acid** obtained by CrO<sub>3</sub> oxidn. of the enzymic product also confirmed the identity of the enzymic **.omega.-hydroxylation** product. This enzymic **hydroxylation** required O<sub>2</sub> and NADPH, but substitution of NADH resulted in nearly half the reaction rate obtained with NADPH. Maximal rates of **.omega.-hydroxylation** occurred at pH 8 and the rate increased in a sigmoidal manner with increasing concns. of palmitic acid. This **.omega.-hydroxylation** was inhibited by the classical **mixed function oxidase** inhibitors such as metal chelators (*o*-phenanthroline, 8-hydroxyquinoline, and **.alpha...alpha.-dipyridyl**), NaN<sub>3</sub>, and thiol reagents (N-ethylmaleimide and p-chloromercuribenzoate). As expected of a hydroxylase, involving cytochrome P450, the present **o**-hydroxylase was inhibited by CO and this enzyme system showed unusually high sensitivity to this inhibition; 10% CO caused inhibition and 30% CO completely inhibited the reaction. Another unusual feature was that the

inhibition caused by any level of CO could not be reversed by light (420-460 nm).

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